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Major gene expression changes and epigenetic remodelling in Nile tilapia muscle after just one generation of domestication

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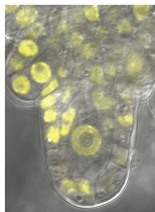
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





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RESEARCH PAPER



Major gene expression changes and epigenetic remodelling in Nile tilapia muscle after just one generation of domestication

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ABSTRACT

The historically recent domestication of fishes has been essential to meet the protein demands of a growing human population. Selection for traits of interest during domestication is a complex process whose epigenetic basis is poorly understood. Cytosine hydroxymethylation is increasingly recognized as an important DNA modification involved in epigenetic regulation. In the present study, we investigated if hydroxymethylation plays a role in fish domestication and demonstrated for the first time at a genome-wide level and single nucleotide resolution that the muscle hydroxymethylome changes after a single generation of Nile tilapia (*Oreochromis niloticus*, Linnaeus) domestication. The overall decrease in hydroxymethylcytosine levels was accompanied by the downregulation of 2015 genes in fish reared in captivity compared to their wild progenitors. In contrast, several myogenic and metabolic genes that can affect growth potential were upregulated. There were 126 differentially hydroxymethylated cytosines between groups, which were not due to genetic variation; they were associated with genes involved in immune-, growth- and neuronal-related pathways. Taken together, our data unveil a new role for DNA hydroxymethylation in epigenetic regulation of fish domestication with impact in aquaculture and implications in artificial selection, environmental adaptation and genome evolution.

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
Domestication; epigenetics;
DNA hydroxymethylation;
muscle growth; *Oreochromis niloticus*

Introduction

For thousands of years, domestication of plants and terrestrial mammals has shaped human evolution. Our civilizations, societies, global demography and diet have been inextricably linked with this process [1]. Historical evidence of fish domestication is recorded much later than that of terrestrial animals. Its first documented instance was when freshwater fishes were attracted to rice paddies and managed alongside them [2]. Domestication is a rather long and complex process, whereby wild animals are not simply kept in captivity, but bred for enhanced traits that are beneficial to humans. Phenotypic, behavioural as well as genetic changes are known to be linked to mutations and shifts in allele frequencies that occur gradually after consecutive generations of selection [3–6]. Nevertheless, phenotypic changes are often detected within a single generation of domestication.

For example, selection for growth can result in an average increase in body weight of 13% per generation in aquatic species of commercial value [7]. Selected lines of Atlantic salmon (*Salmo salar*) demonstrate an increase of 4.6% in feed efficiency ratio per generation compared to their wild counterparts, [8] and weight gain in Nile tilapia can reach up to 14% per generation [9]. Epigenetic mechanisms that promote phenotypic remodelling without involving changes in the nucleotide sequence are expected to be involved in domestication. 5-hydroxymethylcytosine (5hmC) is an epigenetic DNA modification produced through the oxidation of 5-methylcytosine (5mC) by ten-eleven translocation enzymes [10]. Recent evidence pinpoints the importance of 5hmC in epigenetics and its association with active histone marks, histone accessibility and gene regulation [11–13]. Its particular enrichment within gene-bodies; mainly within

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promoters and intragenic regions, as well as its linkage to gene-rich genomic regions suggest that 5hmC is a transcriptionally relevant epigenetic modification [14,15]. Depending on the location, 5hmC enriched regions can be functionally relevant by maintaining an active or inactive chromatin state [14]. Compared to 5mC marks, hydroxymethylation seems to be involved in highly complex mechanisms by activating secondary molecules such as non-coding RNAs. In particular, increased levels of 5hmC at the promoter of microRNA-365-3p, resulted in the regulation of nociceptive behaviours in mice through a potassium channel (Kcnh2), which also pinpoints the accuracy and effectiveness of such modifications to rapidly transfer and regulate environmental signals through the peripheral and central nervous systems [13]. Furthermore, in a state of hypertrophied cardiac muscle cells in mice, 5hmCs are shifting from gene bodies to intergenic regions. Cytosine hydroxymethylation is a pre-activating mark of chromatin remodelling, a process that is crucial for chromatin accessibility [11]. To date, the role of 5hmC in teleosts and their domestication remains unknown.

Here we reveal that the muscle hydroxymethylome changes when the offspring of wild Nile tilapia are reared in captivity for just one generation of domestication. We observed an overall downregulation of immune genes, possibly linked to pathogen exposure and immune cell infiltration in muscle of wild individuals, concomitantly with an upregulation of metabolic and myogenic genes. By focusing on differentially expressed (DE) genes that were associated with differentially hydroxymethylated cytosines (DhmCs), we demonstrate that 5hmC may play a key role in the regulation of cell cycle, metabolism and muscle growth.

Results

The nuclear but not the mitochondrial muscle hydroxymethylome differs between wild Nile tilapia and their offspring reared in captivity

Due to the nature of the reduced representation hydroxymethylation profiling (RRHP) libraries, samples with higher levels of hydroxymethylation produced higher number of hydroxymethylated fragments. Therefore, all libraries were sequenced with equal volumes for proportional sequencing. On average, 69

and 29 M raw reads were obtained per library from the W and D groups, respectively. Approximately 54% of the reads were uniquely mapped to the reference genome, while 14% were multiple mapped (Fig. S1). In total, we covered 1.2 out of 1.8 million possible CCGG sites from both strands across all chromosomes and scaffolds. After removing 5hmC sites having less than 26 counts across 7 or more samples (see Methods), 298,289 CCGG sites were used for further analyses.

Hydroxymethylation was present throughout the whole genome without forming any particular clusters in specific chromosomes (Figure 1(a)). The overall 5hmC pattern across the genome showed increased hydroxymethylation levels in the W group (uniquely mapped CCGG sites; one-tailed Student's t-test; $p < 0.01$). Liquid chromatography/mass spectrometry (LC-MS/MS) confirmed that the global 5hmC levels were significantly higher in the W compared to the D group (Figure 1(b)).

The significant differences in library sizes between the two groups did not reflect similar effects on 5hmC variation, which was higher within the D group (ANOVA; $D_{\text{var}} = 1.54 \times 10^{14}$, $W_{\text{var}} = 1.51 \times 10^{14}$, $F = 9$, $F_{\text{crit}} = 5$, $p = 0.01$ with $\alpha = 0.05$; Table S1), and may be due to relaxed natural selection and phenotypic divergence in captivity conditions [16]. However, the overall variation of individual CCGG sites after filtering was higher in the W group and this variance was mainly associated with large differences between 5hmC sites. The opposite effect was observed in the D group, where the variance was attributed mainly to within-site variation (Table S2). Based on the annotation of all CCGG sites as well as the sites that were hydroxymethylated in our data set, we observed that 5hmCs were relatively depleted within intergenic regions (19%) and enriched in exons (31%) and promoters (33%) (Figure 2).

A total of 13 of the 38 possible mitochondrial CCGG sites were substantially hydroxymethylated (Figure 1(c)). Four of these sites were detected within the *cox1* (cytochrome c oxidase subunit I) gene, which is involved in a large number of processes including metal ion binding, aerobic respiration, and oxidative phosphorylation. Notably, the highest 5hmC levels per CCGG site were observed within the 12 S and 16 S ribosomal RNA genes.

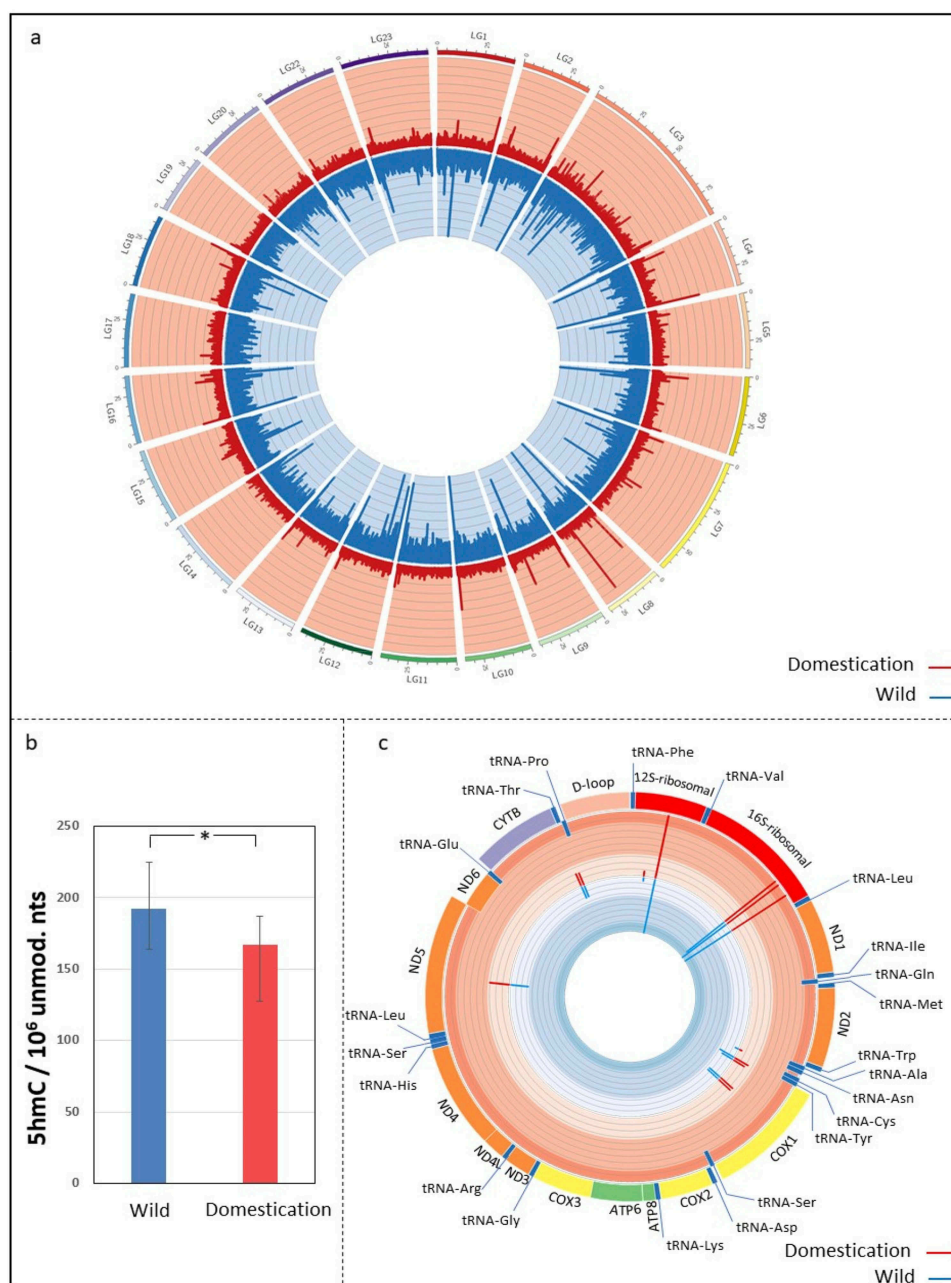


Figure 1. Hydroxymethylation differences between wild Nile tilapia and their offspring reared in captivity. (a), Circular representation of the Nile tilapia nuclear genome (linkage groups LG1-LG23, no scaffolds included for visualization purposes). Only sites with substantial levels of 5hmC were included (298289 CCGG sites). (b), Histogram with the average number of hydroxymethylated cytosines, 5-hm(dC), per 1 million nucleotides, in fast muscle of wild fish compared to their offspring undergoing domestication. Significance was evaluated using the one-tailed Student's t-test ($n = 6$, $p = 0.04$). Standard errors of the means are shown by black bars. (c), Circular representation of the Nile tilapia mitochondrial genome with its gene features in both L and H strands. (a,c), Blue and red peaks (oriented in and out, respectively) represent the hydroxymethylation levels in the groups of wild fish and their offspring undergoing domestication, respectively. The height of each peak is the average count for each group per location ($n = 6$). The 5hmC counts per genomic location and individual, as well as the standard deviation for the W and D groups in panels 1a and 1c can be found in Supplementary Table S9.

Differentially hydroxymethylated genes are involved in immunity, as well as in various intra- and extracellular processes

We identified 126 differentially hydroxymethylated cytosines (DhmCs) between the two groups (p adj.

val. <0.05 ; Figure 3). All sites were hyper-hydroxymethylated in the wild group compared to the first generation of fish reared in captivity. DhmCs were mostly enriched within linkage group 3 but no DhmCs were found in chromosomes LG5,

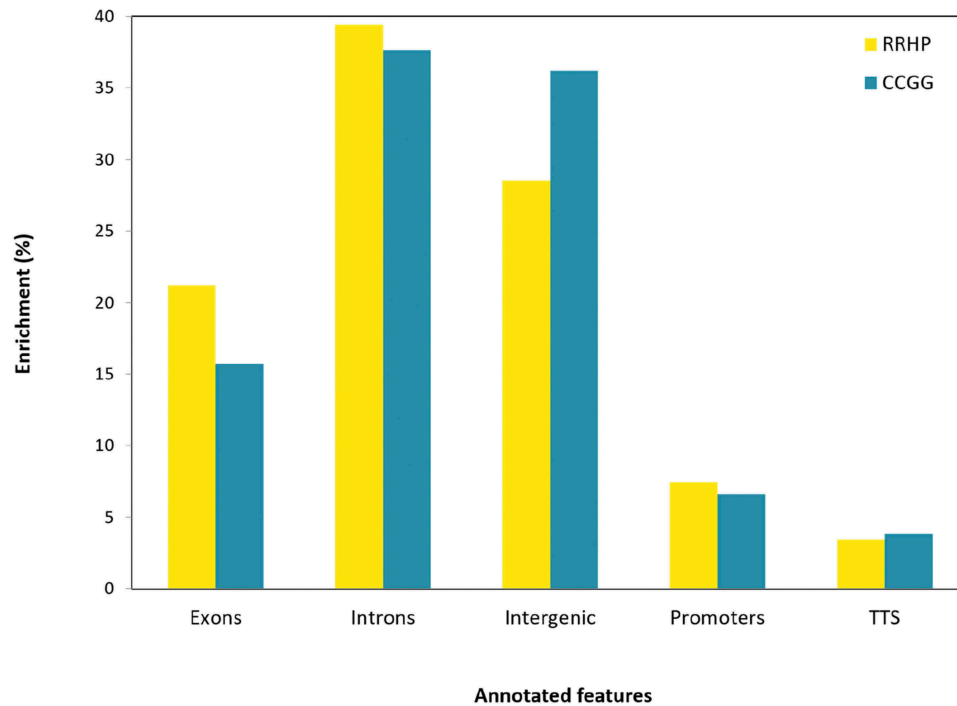


Figure 2. Histogram of gene features (Promoters defined from -1kb to $+100\text{bp}$; TTS defined from -100bp to $+1\text{kb}$). Yellow and turquoise represent the enrichment of sites with substantial levels of 5hmC (RRHP – 295720 sites) and total CCGGs (whole genome – 1791,720 sites) across the Nile tilapia genome, respectively.

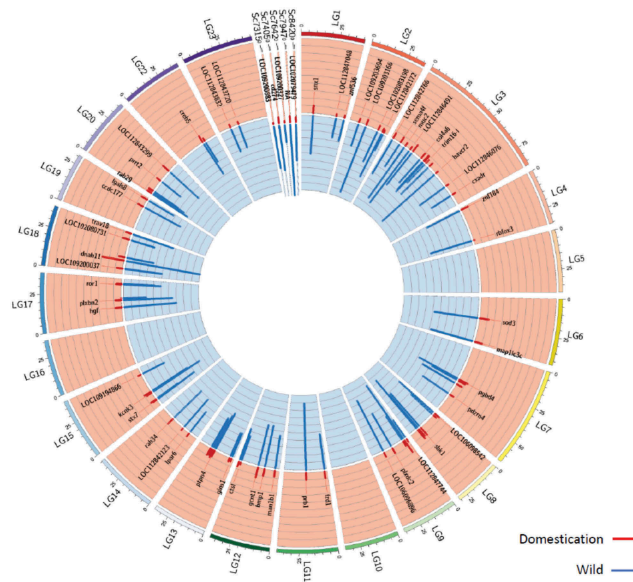


Figure 3. Circular representation of the Nile tilapia nuclear genome (linkage groups LG1-LG23 and scaffolds) showing 126 Dhmc sites. For visualization purposes the numbers in the scaffold symbols 'Sc' represent the last four digits in the accession number starting with 'NW_02032' in the NCBI database (e.g. 'Sc7315' represents scaffold 'NW_020327315'). Gene symbols refer to the closest gene to the reported CCGG site. See Figure 1(a,c) for explanations of peaks. The 5hmC counts per genomic location and individual, as well as the standard deviation for the W and D groups in Figure 3 can be found in Supplementary Table S10.

LG10, LG16 and the mitochondrial genome. Most of these Dhmc sites were located in gene bodies (79) rather than in intergenic regions (47) (Fig. S2). Dhmc sites located in intergenic regions were assigned to genes based on their proximal distance to the transcription start site (TSS) of the corresponding gene. In total, 61 and 34 unique genes were linked to intragenic and intergenic Dhmc marks, respectively. Proportionally to the genomic and filtered CCGG sites, Dhmc sites were enriched within transcription termination sites (genomic-TTS; $p = 0.055$, filtered-TTS; $p = 0.041$, binomial test). The genes with the highest number of Dhmc sites were *myo5b* (myosin Vb) and *kmt2e* (inactive histone-lysine N-methyltransferase 2E) with 4 Dhmc sites, as well as the *lpar6* (lysophosphatidic acid receptor 6), *map1lc3 c* (microtubule-associated protein 1 light chain 3 C) and *extensin* with 3 Dhmc sites (Table S3). Genes that were differentially hydroxymethylated are involved mainly in immune response (*fcgr2b*, *stx7*, *cd274*, *havcr2*, *ctsl* and *sh2d1a*), intracellular processes (*myo5b*, *rab29*, *rab34*, *map1lc3 c*, *elovl4* and *gcnt1*), and core extracellular processes (*bmp1*, *col4a6*, *ctsl*, *hgf*, *muc2*, *plxnc1* and *plxdc2*) (Table S4). These genes are known to play crucial roles in the modulation of response to growth factors and cell growth

inhibition, and they are associated with the canonical pathway of the muscle that is affiliated to structural or functional extracellular matrix proteins ($p = 0.007$). By using the zebrafish annotated features, we identified genes that are involved in axon (GO:0061564; $\text{Log}(p) = -2.67$), neuron (GO:0048666; $\text{Log}(p) = -2.37$) and neuron projection development (GO:0031175; $\text{Log}(p) = -2.16$) as well as the phagosome pathway ($\text{Log}(p) = -2.26$) (Table S5).

A single generation of domestication is associated with substantial changes in the muscle transcriptome

The average size of the rRNA depleted libraries was 19.2 M raw reads for the W samples and 19.3 M raw reads for the D samples (Fig. S3). Of these, 13.1 M and 12.8 M reads were uniquely mapped to the Nile tilapia for the W and D samples, respectively. Of the mapped reads, 3.6 M and 3.8 M reads were mapped to multiple locations in the genome for the W and D samples, respectively.

We found 2,121 genes ($q < 0.05$, $|\log \text{FC}| > 1.5$) that were differentially expressed between the W and D generations. A substantial number of genes (2015) were downregulated in the D group, while only 106 genes were upregulated ($q < 0.05$, $|\log \text{FC}| > 1.5$; Figure 4). A large proportion of the downregulated genes were enriched for immune functions, such as lymphocyte activation ($\log(p) = -24.11$), cytokine production ($\log(p) = -21.6$), myeloid leukocyte activation ($\log(p) = -20.49$), chemotaxis ($\log(p) = -20.26$) and leukocyte migration ($\log(p) = -18.86$), as well as the cytokine-mediated ($\log(p) = -15.68$) and immune response signalling pathways ($\log(p) = -16.98$) (Fig. S4; Table S6). Upregulated genes were involved in metabolic processes such as monocarboxylic acids ($q = 3.43 \times 10^{-5}$), fatty acids ($q = 15 \times 10^{-2}$) and glucose catabolic process to pyruvate ($q = 3.3 \times 10^{-2}$), as well as the glycine, serine and threonine metabolism KEGG pathway ($q = 0.016$). Interestingly, significantly enriched genes within biological functions such as striated muscle tissue development ($q = 3.4 \times 10^{-2}$) and muscle system process ($q = 1.9 \times 10^{-3}$) were also upregulated in the D group, which may indicate

a higher growth potential of fish in captivity (Fig. S5, Table S7).

Hydroxymethylation and gene expression are positively associated

A total of 11 DhmcS were found within 7 DE genes (Table 1; $q < 0.05$, $\log \text{FC} > 1.5$). These 7 genes were hyper-hydroxymethylated and upregulated in the wild group, indicating a positive association between hydroxymethylation and gene expression. The gene that displayed the highest fold change in terms of gene expression was *sh2d1a* (SH2 domain containing protein 1A), which is involved in cellular defence and regulation of immune responses.

The remaining DE genes containing DhmcS play roles in cell survival (*creb5*) and broad neuronal-related effects such as axon guidance and development, neuron projection guidance and development, regulation of axonogenesis and generation of neurons (*dok2*, *plxnc1*, *dnah11* and *myo5b*). A gene essential for the structural maintenance of the extracellular matrix and wound healing of the skin in mammals (*bmp1*) also contained DhmcS and was DE between the two groups. Notably, DhmcS within DE genes were mostly located within introns (8/11), which might affect alternative splicing events or cryptic promoters. Finally, distance-based association between DhmcS and DE genes without the previously mentioned thresholds revealed one additional gene (*fzd1*; $p < 0.05$, $\log \text{FC} = 1.3$) which is involved in a broad range of processes within the Wnt signalling pathway and was differentially hydroxymethylated at a distance of approximately 23 kbp upstream of its transcription start site.

No evidence of population structure between W and D groups

Considering the unknown genetic background of wild Nile tilapia males and their contribution to the D group genotypes, we investigated whether genetic mutations within the restriction sites (CCGG) could potentially explain the observed epigenetic reprogramming during the first generation of domestication. In total, 2,871 single nucleotide polymorphisms (SNPs) were identified among

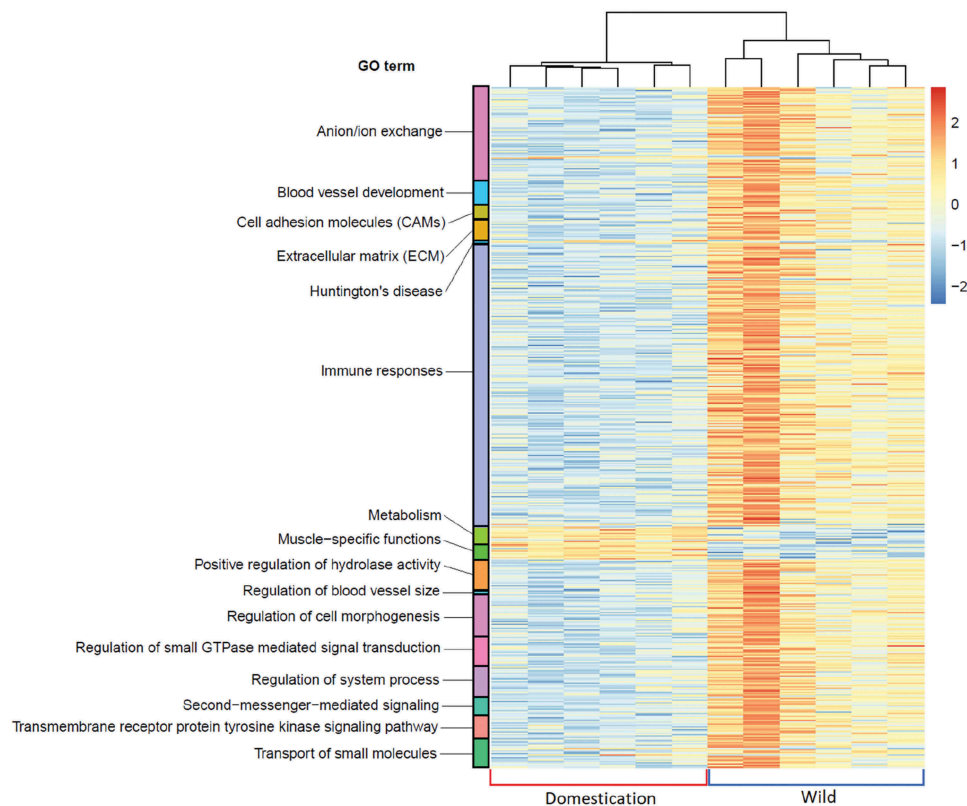


Figure 4. Heatmap showing the differentially expressed genes assigned to their gene ontology (GO) terms between the two clustered groups. Red and blue brackets at the bottom of the heatmap correspond to the 'Domestication' and 'Wild' groups, respectively. The gradient from blue to red colour represents down- to upregulation, respectively. Each GO term is colour-coded as shown at the left side of this heatmap. All related statistics, including significance of each enriched GO term, are shown in tables S5 and S6.

all samples from the same sequencing data that was used to identify 5hmCs. Pairwise SNP distances confirmed that each D-offspring corresponds to each W-parent, considering that the smallest distances were observed between each W-D pair (Fig. S6). The fixation index between the W and D groups was negative ($F_{st} = -0.05$), which confirms our hypothesis that there was no genetic structure between the two Nile tilapia populations, as shown in Figure 5. The within-group nucleotide diversity estimates were similar for both populations.

Discussion

Major leaps in growth rate during the first generation of fish domestication cannot be explained by genetic adaptation alone, such as mutations and selective sweeps that lead to specific shifts in allele frequencies. As shown before, genetic adaptation

to captivity is a relatively slow, species-specific process and can only be detected after several generations [3,4,6,17]. Nevertheless, within a single generation, hatchery-reared fish are selected for traits that are advantageous in captivity (e.g., growth rate, egg size, fecundity, physiological processes associated with smoltification and relaxed anti-predator behaviour) but can be maladaptive in the wild, resulting in reduced reproductive fitness [18,19]. Recent evidence has shown that gene expression patterns can be altered within the first generation of captivity rearing and that these changes can be transferred to the next generation [20]. Indeed, wild growth-hormone transgenic coho salmon (*Oncorhynchus kisutch*) and rainbow trout (*Oncorhynchus mykiss*), appear to have similar modifications in genetic pathways as their counterparts undergoing domestication [21,22]. However, very limited research has been made to uncover the mechanisms underlying these

Table 1. List of genes that were both differentially hydroxymethylated and differentially expressed between wild Nile tilapia (W) and their offspring undergoing domestication (D). P adj.: FDR adjusted p value (q value), log FC: logarithmic fold-change, where negative values represent upregulation in D and positive values represent upregulation in W.

Gene symbol	creb5	myo5b	myo5b	myo5b	myo5b	myo5b	myo5b	myo5b	plxnc1	dok2	dok2	frizzled-1	sh2d1a	dnah11	bmp1
Protein	cyclic AMP-responsive element-binding protein 5	myosin VB	myosin VB	myosin VB	myosin VB	myosin VB	myosin VB	myosin VB	plexin-C1	docking protein 2	docking protein 2	frizzled-1	SH2 domain containing 1A	dynein heavy chain 11 axonemal	bone morphogenetic protein 1
Gene Expression (q value)	0.004	0.009	0.009	0.009	0.009	0.009	0.009	0.009	0.013	0.021	0.021	0.023	0.031	0.036	0.048
Gene Expression (log FC)W vs D	2.3	1.991	1.991	1.991	1.991	1.991	1.991	1.991	2.162	2.022	2.022	1.276	2.688	1.703	1.608
DhmC	3431	4816	4818	3742	3977	-2695	7015	7017	8999	7153	4250				
(distance_to_TSS)															
DhmC (Annotation)	intron 3 of 10	intron 1 of 38	intron 1 of 38	intron 1 of 38	intron 1 of 38	intron 1 of 38	intron 3 of 6	intron 3 of 6	TTS	Intergenic	exon 38 of 40	TTS	intron 1 of 19		
DhmC (log FC)W vs D	1.457	6.242	6.233	5.655	6.421	6.534	5.660	6.414	0.008	0.008	0.002	1.075	6.481		
DhmC (P adj.)	0.008	0.008	0.008	0.008	0.008	0.008	0.018	0.008	0.008	0.008	0.002	0.030	0.009		
Strand	-	-	+	-	-	+	-	-	-	-	+	+	+		
Position	25495940	36567964	36567962	36569038	36568803	34318460	37756187	37756189	2292991	31466924	12201282	13634977			
Chromosome	NC_031985.2	NC_031972.2	NC_031972.2	NC_031972.2	NC_031972.2	NC_031981.2	NC_031977.2	NC_031977.2	NC_031976.2	NC_031966.2	NC_031982.2	NC_031977.2			

changes. In the present study, 2,121 DE genes were identified between wild Nile tilapia and their first-generation offspring. A rapid adaptation to captivity conditions was evident at the molecular level, with most genes being downregulated in the group undergoing domestication. In a recent study [20], 723 genes were reported to be DE between wild-born and first generation hatchery-reared offspring Steelhead trout (*Oncorhynchus mykiss*). Unlike our findings in Nile tilapia, most of these genes were upregulated in hatchery-reared fish and only the muscle-related troponin I gene (*tnni1*) was found to be upregulated in both steelhead trout and Nile tilapia reared in captivity. This might be an indication of either species- or tissue-specificity (whole fry vs fast muscle) or even differences in environmental factors (e.g., rearing conditions, temperature, dissolved oxygen and pH) that contribute to the expression of these genes.

A total of 316 genes directly associated with the immune response were upregulated in wild Nile tilapia compared to their counterparts undergoing domestication. These results demonstrate the challenging and variable environmental factors affecting wild populations, namely, high water temperatures during the summer months (up to 35°C at river banks and shallows), exposure to viruses, bacteria and parasites, as well as predators and fishing pressure compared to the predator- and pathogen-free conditions of our recirculating aquaculture system. For example, one of the major physiological differences between the W and D group was the obvious presence of endoparasites in the W group (supplementary video 1), which could justify the upregulation of several genes such as interleukins and their receptors (*il21 r*, *il12rb2*, *il12rb1*, *il16*, *il2rb*, *il7 r*, *il18rap*, *il1rap*, *il17 f*, *il1r2*, *il4l1*, *il1b*, *il31ra*) and other genes previously reported to be upregulated as a response to parasitic load in vertebrates [23] (e.g., *irf4*, *irf1*, *thbs1b*, *socs3*, *junb*, *prf1*, *ccr9*, *itga5* and *slc3a2*). Based on the tissue enrichment analysis and manual inspection of expression patterns of these genes in humans (p adj. val. $\approx 10^{-7}$; Fig. S7), it is plausible that wild fish had a higher degree of immune cell infiltration in muscle tissue than their offspring reared in captivity, even though this could not be confirmed by histological analysis due to the unavailability of necessary chemicals and equipment in the field.

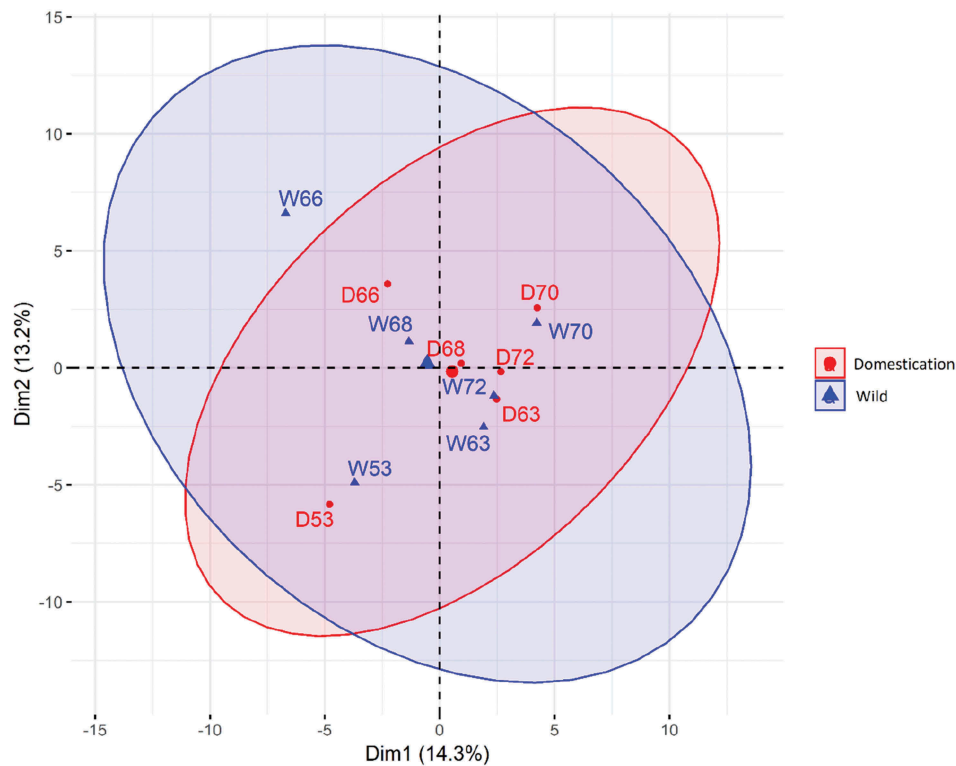


Figure 5. Principal component analysis (PCA) showing the genetic diversity of the 6 wild and 6 first-generation offspring based on their SNPs. Blue triangles and red circles represent the wild fish (W) and their counterparts in the first generation of domestication (D), respectively.

It is noteworthy that T and B cell activation, differentiation, and function is mediated by ten-eleven translocation enzymes and the DNA oxidation and demethylation pathways [24]. We found that *tet3* was upregulated in the W group ($p < 0.05$, $\log FC = 1.17$), which suggests an overall demethylation process within both muscle and immune related cells, with major implications in gene transcription. These findings demonstrate the complexity of epigenetic mechanisms and their roles in gene regulation, cell defence and growth.

The overall muscle composition changes with age and the different cell types present in muscle have distinct DNA methylation patterns [25]. As the exact age of the wild Nile tilapia could not be determined by otolith analysis, one cannot exclude the possibility that the observed upregulation of *tet3* in the W group was at least partly due to age differences. Nevertheless, the local fishermen that assisted us in Egypt know that these fish were approximately 4–5 months old due to the fishing pressure imposed by the gill nets and traps used. The changes in gene expression and 5hmC levels identified between wild fish

and their progeny undergoing domestication are most likely due to environmental differences, since the latter were also 5 months old at the time of sampling. Stress-related epigenetic changes may also play an important role in immunocompetence and acute stress can increase the transcription of immune-related genes [26]. Even though the authors have taken precautions towards reducing any bias in the experimental design, the collection of fertilized eggs from the mouth of the W females without anaesthesia represented an additional acute stress that could potentially have influenced the epigenome.

Crowded conditions can potentially result in skin abrasions and hypoxia [27] and rearing density plays an important role towards the adaptation to captivity [20]. The rearing density in our recirculating aquaculture system was higher than in the wild but we found few DE genes related to crowding conditions. Specifically, we identified only two genes, *grn* (granulin) and *phb2* (prohibitin-2) that were upregulated in fish reared in captivity and that were related to wound healing and response to hypoxia, respectively.

Studies on commercial fish [28–31] and model organisms [32], have shown that captivity rearing usually offers the appropriate conditions for increased growth rates. Phenotypic selection has led to the development of the genetically improved farmed tilapia strain, which gradually gained up to 15% in body mass per generation within the first six generations of domestication [33]. Considering the impact of environmental factors on the epigenome and the overall health status during the transition from the wild to an enclosed rearing environment, it is plausible that the first signs of growth-gain are strongly associated with a lower pathogen load in captivity. This notion is further supported by the fact that the immune system is closely associated with the muscle development, since chronic inflammation can lead to muscle damage and fibrosis [34]. It is likely that other mechanisms directly affect muscle growth during the first generation of domestication. We found two genes (*myo5b* and *creb5*) that are both DE and contain DhmcCs between wild and first-generation captive fish. These genes are involved in the PI3K-Akt signalling pathway, which is one of the most important intracellular pathways in muscle that regulate cell cycle, metabolism, proliferation and growth. Myosin Vb targets and activates directly the phosphatase and tensin homolog (PTEN) [35], which in turn dephosphorylates PIP₃ (phosphatidylinositol 3,4,5 – trisphosphate) and prevents cells from growing. Both *myo5b* (LogFC = 1.991; q = 0.009) and *pten* (LogFC = 1.303; q = 0.043) were upregulated in wild individuals, suggesting a possible epigenetic mechanism that regulates muscle hypertrophy. Additionally, *myo5b* is directly involved in insulin-dependent glucose transporter 4 (GLUT4) translocation in muscle cells and uses *rab8a* and *rab13* GTPase-activating proteins that require Akt signalling and activation [36,37]. Apart from its involvement within the PI3K-Akt signalling pathway, *myo5b* plays various roles in neuronal processes. It recycles the endosome delivery into hippocampal neuron spines in dendrites, targets and activates PTEN, which in turn regulates the soma size, and promotes axonal development in conjunction with the Ras-related protein *rab10* [38]. It would not be surprising if the epigenetic regulation of neuronal development was associated with muscle development, considering that neurons are an integral part of the skeletal muscle. *Creb5* codes for

a protein that is a substrate for the Akt/PKB and is involved in cell survival. Among other pathways, *creb5* is also involved in insulin resistance and gluconeogenesis in the muscle, which occurs during periods of starvation and low-carbohydrate diets [39]. *Myo5b* and *creb5* seem to be good candidates for further investigation into their epigenetic regulation during domestication. Overall, genes that were upregulated in the D group were mostly related to metabolic processes (43/106). We also identified genes involved in myosin binding and actomyosin structure organization (*actc1*), regulation of skeletal muscle contraction (*casq2*, *eno1*, *mb*) and skeletal muscle development (*srpk3*), response to and regulation of glucose import (*rtn2*, *mef2a*) as well as calcium release channels and insulin secretion (*fkbp1b*). Interestingly, *srpk3* (serine/arginine-rich specific kinase 3), is a well conserved gene among mammalian species and it is closely related with myocyte-specific enhancers that promote muscle growth in the skeletal muscle tissue [40]. These results coupled with the dietary effects in captivity, such as high-quality diet and regular feeding *ad libitum*, suggest an increase in metabolic capacity that leads to a higher growth rate from the very first generation of fish undergoing domestication. It is plausible that this effect is partly due to nutritional programming. Previous studies in zebrafish (*Danio rerio*) have shown that parental nutritional programming through increased dietary intake of arachidonic acid or deficiency in 1-C nutrients results in DNA methylation changes and epigenetic regulation of gene expression in the liver of the offspring [41,42].

The total number of 5hmCs in our reduced representation of the Nile tilapia genome was substantially high, covering ubiquitously the nuclear genome as well as identifying 5hmCs within the mitochondrial genome. High levels of 5hmC within the 12 S and 16 S ribosomal RNA might indicate that epigenetics could have an effect in the translational dynamics of mitochondrial mRNAs. Although studies have shown 5hmCs to be present in mammalian mitochondria [43,44], there are no reports of 5hmCs at single nucleotide resolution in the mitochondria of teleosts. Here we provide the first evidence of 5hmC marks across the mitochondrial genome of Nile tilapia, which also implies the existence or relocation of CpG demethylation mechanisms within the mitochondria. Strikingly, the large differences of the

nuclear 5hmC levels between wild and first-generation fish, do not reflect the average 5hmC levels within the mitochondria. This suggests that the two demethylation mechanisms are effectively separated. Similarly to our study, 5hmCs in the muscle of zebrafish occur mainly in bodies of genes, some of which have functions with interesting implications in metabolism, development and other muscle-specific functions [45].

We observed higher hydroxymethylation levels in the W group, which resulted to the identification of 126 DhmcS. A previous study that compared the DNA methylation profiles of hatchery-reared and wild coho salmon (*O. kisutch*), identified 37 differentially methylated regions (DMRs) that are likely linked with the reduction in fitness of the hatchery-reared fish [46]. These DMRs were found within 52 unique transcripts, but none of them appeared in our Dhmc data. Possible explanations for these differences are species-specificity (smoltification related processes that are not present in Nile tilapia), ontogeny stage (fry vs adults), and 5mC/5hmC-specificity (different epigenetic modifications may control different pathways). There were two genes in common between our DE genes and these 52 transcripts, namely ankyrin-1 (*ank1*), which is involved in cytoskeleton organization, and *phb2*, which is involved in a large number of processes, such as cellular response to hypoxia and response to wounding. *Ank1* was upregulated in W fish, whereas *phb2* was upregulated in the D group, which indicates a common regulation of these genes among fish species during the early stages of domestication. This finding possibly indicates that similar stressors might affect the wild populations of fish with major impact on their fitness and growth potential. However, our data indicate that most molecular networks regulated during the early stages of domestication appear to be species-specific.

According to a recent study, DNA methylation of the first intron has a negative effect on gene expression levels across several species and tissues [47]. In our case, the first intron of *myo5b* is highly hydroxymethylated in the W group (4 DhmcS, logFC = 5.69, q val = 0.0085) compared to the D group. In addition, we observe a positive correlation with its transcript levels (logFC = 1.99, q val = 0.009). Additionally, a single Dhmc was found within the first intron of *bmp1* (logFC = 6.48,

q val = 0.008) and it was also upregulated in the W group (logFC = 1.6, q val = 0.048). These results indicate the importance of distinguishing 5mC from 5hmC calls. Unlike our RRHP approach, other commonly used methods such as reduced representation bisulphite sequencing (RRBS) are unable to do so, which limits the conclusions from those studies.

While genetic variation alone did not explain the large differences in 5hmC patterns between the W and D groups, it is worth mentioning that the SNP coverage was limited due to the nature of the RRHP libraries. A broader genome-wide approach could potentially identify additional SNPs between the two groups. Furthermore, hydroxymethylation levels at the beginning of the embryonic development in fish are very low [45,48], therefore epigenetic reprogramming is likely occurring as a result of environmental conditions rather than of maternal effects.

Conclusion

The process of domestication affects animals in ways that still remain unknown. Both genetic and epigenetic changes can alter the phenotype within a single generation. Here, we demonstrated for the first time that DNA hydroxymethylation in muscle differs significantly between wild fish and their progeny reared in captivity. We further identified DhmcS mostly within gene bodies, which might indicate their functional role in epigenetic regulation. These DhmcS were associated mainly with immune-, neuronal- and growth-related processes. Concurrently, thousands of genes were differentially expressed between groups, with upregulation of immune related genes in the wild and the upregulation of metabolic and muscle-specific genes in the first generation of fish undergoing domestication. Notably, genes that were found to be both differentially hydroxymethylated and differentially expressed had a consistently positive correlation. Another remarkable finding was the hyperhydroxymethylation of the first intron of *myo5b* and its connection with the upregulated *pten* gene in the wild group. *Pten* is involved in the PI3K-Akt signalling pathway and muscle atrophy. Further investigation into the PI3K-Akt pathway could reveal one plausible way for rapid regulation of growth and muscle hypertrophy in fish during the first stages of their domestication. Lastly, we

found no evidence for genetic differentiation between the two groups, showing that the observed hydroxymethylation changes within the first generation in captivity have an epigenetic rather than genetic basis. Understanding the epigenetic mechanisms, namely hydroxymethylation, involved in early domestication is essential for aquaculture diversification (farming of new species), sustainable production, and wild fish stock management practices.

Materials and methods

Ethics approval

This study was approved by the Nord University (Bodø, Norway) ethics board and all procedures involving animals were performed according to the instructions of the Norwegian Animal Research Authority (FOTS ID 1042).

Experimental design

A total of 6 wild (W) mouthbrooding Nile tilapia females were captured using traps and gill nets along the river Nile in Luxor, Egypt within a 500 m radius from the following coordinates (latitude: 25.6655° N; longitude: 32.6186° E). After collecting approximately 180 fertilized eggs from each female by gently holding their head in containers filled with river water, the fish were placed in 25 L tanks, transferred from the boats to the sampling location and sacrificed by immersion in a clove oil (Sigma Aldrich, USA) emulsion containing 12 ml of a clove oil pre-mix in 96% ethanol (1:10 v/v) in 10 L of water [49].

A 0.5 cm thick cross-section was taken at 0.7 standard length, just anterior to the urogenital opening. Fast (white) skeletal muscle was carefully dissected from the left dorsal quadrant of the section and preserved in DNA/RNA Shield (Zymo Research, USA) at 4°C. Sixteen days post-sampling, all tissues were stored at -20°C. Each batch of eggs was disinfected with hydrogen peroxide (1 g H₂O₂/L for 10 minutes) and placed in egg rockers (Cobalt Aquatics, USA) installed in a 60 L tank with UV-treated water, containing 5‰ NaCl (w/v). Approximately, 85% of the eggs hatched within 4 days at 28°C. At 9 days post-fertilization, the larvae

were placed in fish transport bags filled with UV-treated and 100% oxygen saturated water, and were air freighted to Nord University's research station (Bodø, Norway). Transportation of the larvae lasted approximately 18 hours and their survival rate exceeded 95%. They were reared for 5 months at a maximum density of 27 fish/m³ in a freshwater recirculating aquaculture system (pH = 7.6, DO = 100%, temperature = 28°C and photoperiod LD 13:11) and were fed *ad libitum* with 0.15–0.8 mm Amber Neptun pellets (Skretting, Norway). The fish reared in captivity represent the first generation of fish (D) undergoing domestication. Each D group was kept separate in different tanks based on their ancestry. Fast muscle from 6 D females was sampled and stored using the same procedure as above. W and D samples were genetically paired (i.e., each D individual corresponded to a W progenitor) and at approximately the same age and ontogeny stage (i.e., puberty). All sampling conditions at the research facilities were simulated to match the sampling of the wild fish in Egypt. To compensate for the starvation period of wild mouthbrooding females, D individuals were starved for 48 hours prior to sampling. All fish were sampled at approximately the same time of the day (7–10 am). Both W and D fish were removed from their environment (Nile river for W and rearing 1500 L tanks for D) and kept for a maximum period of 30 minutes in a 25 L tank with anaesthetic before sampling. All sampling procedures were performed in an air-conditioned room at 25°C and oxygen was provided by an air pump. Additional information on the animals, such as total and standard lengths, weight and general observations at the time of sampling can be found in supplementary material (Table S8). Aliquots of the same fast muscle sample from each individual were used for DNA and RNA isolation.

Global 5hmC quantification by LC-MS/MS

Genomic DNA from fast muscle was isolated from all 12 individuals (6 D and 6 W) using the Quick-DNA miniprep Plus kit (Zymo Research). The concentration of DNA was determined using the Invitrogen Qubit 3.0 fluorometer (ThermoFisher Scientific, USA). Quality and integrity of the DNA was assessed using the NanoDrop 1000 Spectrophotometer (ThermoFisher Scientific) and the TapeStation 2200

(Agilent Technologies, USA), with absorbance ratios $260/280 > 1.93$, $260/230 > 1.89$ and minimum DNA integrity number (DIN) of 8.4, respectively. For the global quantification of 5hmC, the internal standard d3-5-hydroxymethyl-2'-deoxycytidine (Toronto Research Chemicals, Canada) was added to the DNA samples. Subsequently, they were hydrolysed completely to deoxynucleosides by 20 U benzonase (Santa Cruz Biotech, USA), 0.2 U nuclease P1, and 0.1 U alkaline phosphatase (Sigma Aldrich) in 10 mM ammonium acetate pH 6.0 and 1 mM magnesium chloride at 40°C for 1 hour. Three volumes of acetonitrile were added to the mix before their centrifugation (16,000 g, 30 min, 4°C). The supernatants were dried and dissolved in 50 µl water for LC-MS/MS analysis of modified and unmodified deoxyribonucleosides. Chromatographic separation was performed using an Agilent 1290 Infinity II UHPLC system with an ZORBAX RRHD Eclipse Plus C18 150×2.1 mm ID (1.8 µm) column protected with an ZORBAX RRHD Eclipse Plus C18 5×2.1 mm ID (1.8 µm) guard column (Agilent Technologies). The mobile phase consisted of water and methanol (0.1% formic acid was added to both phases). For the initial run of 5-hm(dC), the flow rate was set at 0.15 ml/min with 5% methanol for 30 seconds, which was adjusted to a gradient of 5-15% methanol for the next 4 minutes. The flow rate was increased to 0.25 ml/min in a methanol gradient of 15-90% for 3 minutes, and finally re-equilibration was performed at 5% methanol for 4 minutes. A portion of each sample was diluted for the analysis of unmodified deoxynucleosides. Unmodified deoxynucleosides were chromatographed isocratically with 20% methanol. Mass spectrometric detection was performed using an Agilent 6495 Triple Quadrupole system operating in positive electrospray ionization mode, monitoring the mass transitions 261.1/145.1 (d3-5-hm(dC)), 258.1/142.1 (5-hm(dC)), 252.1/136.1 (dA), 228.1/112.1 (dC), 268.1/152.1 (dG), and 243.1/127.1 (dT).

RRHP library preparation and sequencing

The preparation of the reduced representation hydroxymethylation profiling (RRHP) libraries was carried out according to the manufacturer's protocol (Zymo Research). Fast muscle stored in DNA/RNA Shield (Zymo Research) was homogenized using ZR bashing beads in lysis tubes (Zymo

Research) on a Precellys 24 homogenizer (Bertin Instruments, France) at 5000 rpm for 2 cycles of 20 seconds. DNA was extracted using the Quick-DNA miniprep kit (Zymo Research) with a 25 µg binding capacity. After digestion with the restriction enzyme MspI at 37°C for 8 hours, the DNA was washed using the Zymo-Spin IC columns provided in the RRHP 5-hmC library prep kit. Specific adapters that re-create the MspI cutting site were introduced. After library extension, all fragments were glucosylated at 37°C for 2 hours and washed as recommended. During this step, all 5hmCs at the starting CCGG junction of each fragment were glucosylated and protected from the second MspI digestion. At the end of this procedure, only fragments with hydroxymethylated CCGG junctions remain adapterized. Double indices were introduced after size selection according to Illumina's instructions [50] for a 6-plex sequencing design. To minimize bias during library preparation, all samples were handled simultaneously, as suggested in the literature [51,52]. In total, 6 D, 6 W and 2 positive control libraries were sequenced. The positive controls were 5hmC independent-libraries and contained all possible CCGG sites across the genome. Thus, we were able to identify biases in fragment amplification within our count-based matrices. All samples were sequenced at the Norwegian Sequencing Centre with equal volumes in three lanes on a HiSeq4000 (Illumina, USA) to obtain 150 bp paired-end reads. The positive controls were sequenced at 1/3 volume in all three lanes to account for possible lane effects. Due to the low complexity of the libraries (CCGG) at the beginning of each read, 30% PhiX control v3 (Illumina) was added to each lane.

Bioinformatic analyses

Adapters and N bases from the 3' end of each read were trimmed using trim_galore v0.4.4 (Babraham Bioinformatics) [53]. Clean reads were aligned to the reference genome *O. niloticus*_UMD_NMBU [54] with Bowtie v0.12.8 [55] using the parameters -best, -v 1, -n 1 and -m 3. These options compromised alignment rates for robustness and reliability, as they minimized multiple aligned reads. From each SAM alignment file, only reads starting with CCGG from

both strands were extracted and their coordinates (chromosome and position) were used as input into R [56]. A count matrix was created for each sample and afterwards all samples were concatenated into one count matrix. With the aim of avoiding false discovery of 5hmCs, mainly due to within-group variation, we applied double filtering and stringent parameters. At first, a filter was applied in order to remove 5hmCs with low number of counts. All 5hmCs were removed from the matrix when 7 out of 12 samples had below 1 count. This filter not only excluded hydroxymethylated cytosines with low counts but it increased the overall median value and ensured that 5hmC on/off switches remained in the data set. After calculating the median, we also removed 5hmC sites when 7 or more samples had counts lower than the median value of the matrix. In order to perform a comparison of the two groups, our design handled the six different families as blocks while the W and D samples were considered as two separate groups. We then used the R package limma to perform a duplicate correlation based on their counts and the RRHP bioinformatics was performed similarly to gene expression (RNA-seq) rather than methylation (RRBS) data analysis. The Benjamini-Hochberg procedure was used to calculate adjusted p-values for each 5hmC and single cytosines with an adjusted p value ≤ 0.05 were counted as DhmcCs between the W and D groups. The annotation of all significant positions was performed using the software HOMER, annotatePeaks.pl [57] and the NCBI *Oreochromis niloticus* annotation release 104. The association of differentially hydroxymethylated genes with their corresponding gene ontology terms was performed with Metascape [58]. Two separate gene ontology analyses for Nile tilapia genes were performed based on known functions for their annotated (i) human (*Homo sapiens*) and (ii) zebrafish (*Danio rerio*) orthologs.

RNA library preparation, sequencing and analysis

Fast muscle in DNA/RNA Shield was homogenized as described above. Total RNA from the same 12 individuals was isolated using the Quick-RNA miniprep plus kit (Zymo Research) according to the manufacturer's protocol and recommendations for samples stored in DNA/RNA Shield. Ribosomal RNA was removed using the Ribo-Zero Gold rRNA Removal

kit (Illumina). RNA concentration and integrity were assessed using the RNA High-sensitivity tape on the TapeStation 2200 (Agilent Technologies). RNA-seq libraries were prepared using the NEBNext Ultra II Directional RNA Library Prep Kit for Illumina (NEB, USA), according to the manufacturer's instructions. A unique barcode was assigned to each sample and all libraries were pooled at equimolar concentrations. They were then sequenced at the Norwegian Sequencing Centre on a single HiSeq 4000 lane (Illumina), yielding 480 million 150bp paired-end reads. The raw sequencing files were trimmed using trimmomatic v0.33 [59] with the following parameters: *PE -phred33, LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:36*. The trimmed reads were aligned to the Nile tilapia genome (NCBI assembly GCA_001858045.3) using STAR [60] with default settings and paired-end mode. The mapped reads were used as input to *htseq-count* [61] to calculate count numbers for each gene using the gff3 genome annotation from the above mentioned genome assembly. Differentially expressed genes were determined using the edgeR package in R. The differentially expression analysis required that each gene should be expressed with at least 1 count per million in more than 25% of the samples. The p-values for the differentially expressed genes were adjusted for multiple testing using the Benjamini-Hochberg correction [62].

Based on the gene ontology and annotated human orthologs, a high number of immune-related genes in the muscle of wild fish was observed. To better understand the role of these genes in the muscle we manually inspected tissue-specific patterns in human tissues. The analysis was performed using the GeneAtlas U133A gcrma dataset [63] and a plot for tissue-specific gene expression was created using the TissueEnrich tool [64].

Genetic diversity using SNPs from the RRHP sequencing

The RRHP method allows extraction of SNP information across the genome using the same sequencing data [52]. After quality control and adapter trimming, the resulting fastq files were sorted in directories as recommended by the CFSAN SNP Pipeline [65]. The output SNP distance matrix was used to plot distance-based radars among samples. The resulting vcf

files were then used as input for the R package PopGenome with default parameters to calculate the fixation index (Fst) as well as the nucleotide diversity within and between the two groups. A principal component analysis (PCA) was performed using the R package *adegenet* and plotted using the R package *factoextra*.

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Authors' contributions

IK carried out the sampling, wet lab and bioinformatic analysis, interpreted the results and wrote the article. PS contributed significantly to the bioinformatic analysis of the RRHP data set, the interpretation of the results and revised the article. RM contributed significantly to the bioinformatic analysis of the RNA-Seq data set and revised the article. AVN contributed significantly in the wet lab and revised the article. DR contributed significantly to the bioinformatic analysis of the SNP data set. JMOF conceived the study, carried out the sampling, provided reagents and consumables, and contributed significantly to the interpretation of the results and the article revision.

Code availability

The scripts used for transcriptomic and hydroxymethylation analyses are available from the corresponding author upon request.

Disclosure statement

The authors declare that they have no competing interests.

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Data availability statement

The RRHP dataset generated and analysed during the current study is available in the SRA (NCBI) repository, under the accession number PRJNA559277. The RNA-seq dataset generated and analysed during the current study is available in the GEO (NCBI) repository, under the accession number GSE135811.

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